



Protocol DNA Isolation from Tissue & Cells

by nexttec[™] 1^{-Step}

- nexttec™ cleanPlate96 -

Cat. No. 10N.901

Cat. No. 10N.902

Cat. No. 10N.904

Cat. No. 10N.924

Version 1.0

For research only

Principle

nexttec[™]1^{-Step} is the easiest handling and fasted DNA purification system containing a <u>single</u> buffer system and a <u>one-step</u> DNA purification after lysis.

Proteins, detergents and low molecular weight compounds are retained by the nexttec[™] sorbent. DNA passes through the nexttec[™] cleanPlate96 during a short, <u>one-step</u> purification procedure.

The obtained DNA is suitable for all common enzymatic reactions (restriction digests, real-time PCR, PCR, genotyping etc.).

Kit contents

The kit contains all necessary reagents for lysis and subsequent DNA purification.

Component	Art.No. 10N.901	Art.No. 10N.902	Art.No. 10N.904	Art.No. 10N.924
Buffer G	15 ml	42 ml	65 ml	400 ml
Proteinase K	1.5 ml	3 ml	4.5 ml	27 ml
Prep Solution	40 ml	100 ml	150 ml	2x 450 ml
DTT (optional for special protocols)	0.5 ml	0.5 ml	1 ml	5 ml
nexttec [™] cleanPlates96	1	2	4	24
nexttec [™] deep-well plates	3	6	12	72
Sealing tapes	3	6	12	72
Alu sealing tapes	2	4	8	48

nexttec[™] service

For extending the application range to samples, which are difficult to lyse by the standard procedure, it is recommended to include optional components in the lysis buffer, optimizing the lysis time or using different lysis volumes (e.g. robotic applications).

For detailed information, please get in contact with service@nexttec.biz.

Storage Conditions

During shipment all kit components are stable at room temperature. After arrival, **Proteinase** K, Buffer G and Prep Solution must be stored at +2°C to +8°C. After first opening freeze DTT at -18°C to -20°C.

nexttec[™]cleanPlates96 can be stored at room temperature (+20°C to +25°C). If properly stored, see expiration date for the stability of the kit.

Safety Information

Proteinase K Xn R 36/37/38 R42/43; S 23-24-26 36/37

DTT Xn R 22-36/37/38 S 26-36

Risk Phrases

R 22 Harmful if swallowed

R 36/37/38: Irritating to eyes, respiratory system and skin.

R 42/43: May cause sensitisation by inhalation and skin contact.

Safety Phrases

S 23: Do not breathe Gas/fumes/vapour/spray.

S 24: Avoid contact with skin.

S 26: In case of contact with eyes, rinse immediately with plenty of water and

seek medical advice.

S 36/37: Wear suitable protective clothing and gloves.

When working with chemicals, always wear a suitable lab coat, disposable gloves and protective goggles. For more information please consult the appropriate material safety data sheets (MSDS).

Before starting

Equilibrate nexttec[™] cleanPlates96

E1	Add 350 µl Prep Solution onto each well of a nexttec[™] cleanPlate96 . Incubate for at least 5 min at room temperature and centrifuge at 350x g for 1 min or apply vacuum for 30 to 60 sec to remove excess buffer.
E2	Discard the deep-well plate. Place the nexttec TM cleanPlate96 onto a new deep-well plate. Use equilibrated nexttec TM cleanPlates96 or store closed at +2°C to +8°C and use within one week.

Preheat an incubator to 56°C

Protocol

<u>Lysis</u>	
L1	Transfer tissues (5-20 mg fresh weight) or cells (1-2 x 10 ⁶) to a deep-well plate.
	Centrifuge cell suspensions (2,000x g, 10 min), remove and discard the super-
	natant.
L2	Add 140 µl Buffer G, 10 µl Proteinase K and 1.5 µl DTT* to each sample.
L2	Incubate sample with shaking (56°C, 200 rpm, 30 min to overnight).
*For Pr	e-Mixes see Technical Section.
Purifica	ation of DNA
	Transfer 100 μl of the lysates to an equilibrated nexttec™ cleanPlate96.
Р	Incubate for 3 min at room temperature.
	Centrifuge at 700x g for 1 min or apply vacuum for 1 min .
	The eluate contains the purified DNA!!
Notes	<u> </u>

<u>notes:</u>			

Technical Section

Preparation of Lysis buffers (Pre-Mixes)

LG	Lysis Buffer LG:	1 sample	1 plate	2 plates	3 plates	4 plates	
	Buffer G	140 µl	15 ml	31 ml	46 ml	62 ml	
	Proteinase K	10 μΙ	1.1 ml	2.2ml	3.3 ml	4.4 ml	
	Mix by vortexing. Add 150 μl of Buffer LG to each sample (L 2). The Lysis Buffer LG is stable for 1 working day, if stored at +2°C to +8°C .						

Determination of DNA concentration in nexttec[™] 1^{-Step} DNA preparations

We recommend determining the DNA concentration:

- Using the fluorescent dye Picogreen® or similar
- Comparing the fluorescence intensity of DNA bands of unknown concentration with standards, e.g. in ethidium bromide stained agarose gels.

Please notice:

The use of absorption measurement at 260 nm (A_{260}) in a spectrophotometer (e.g. NanoDrop[®]) for determination of DNA concentration <u>is system related</u> not recommended. For details and possible workarounds for your specific application please contact: **service@nexttec.biz**.

Centrifugation

For centrifugation at 350x g or 700x g use the settings for relative centrifugal force (RCF) of your centrifuge. Alternatively measure the distance of the nexttecTM cleanPlate96 to the centre of your rotor and calculate the necessary rotations per minute (e.g. rpm = 299.07 x $\sqrt{350/r}$; r=radius in cm)

• Vacuum Application

Assemble the vacuum manifold as described in the manual. The flow rate should be between 15 L/min and 150 L/min.

A regulation of vacuum is not necessary.

Product Use Restriction

nexttecTM 1^{-Step} DNA Isolation Kit components were developed, designed and sold **for research purposes only**. They are suitable for in vitro uses only. No claim or representation is intended for use to identify any specific organism or of clinical use.

It is the responsibility of the user to verify the use of the nexttec[™] 1^{-Step} DNA Isolation Kit for a specific application as the performance characteristic of this kit has not been verified to a specific organism.

Troubleshooting, FAQ and Special Applications

Product claims are subject to change. Therefore, please, visit our website or contact our technical service team for troubleshooting guide, up-to-date protocols and latest applications on nexttecTM 1^{-Step} products.



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